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Expression of Actin-interacting Protein 1 Suppresses Impaired Chemotaxis of *Dictyostelium* Cells Lacking the Na⁺-H⁺ Exchanger NHE1

Chang-Hoon Choi,* Hitesh Patel,[†] and Diane L. Barber*

*Department of Cell and Tissue Biology, University of California, San Francisco, San Francisco, CA 94143; and [†]University of Edinburgh, ECRC-Cancer Biology, Edinburgh EH4 2XR, United Kingdom

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Increased intracellular pH is an evolutionarily conserved signal necessary for directed cell migration. We reported previously that in *Dictyostelium* cells lacking H⁺ efflux by a Na⁺-H⁺ exchanger (NHE; *Ddnhe1*[−]), chemotaxis is impaired and the assembly of filamentous actin (F-actin) is attenuated. We now describe a modifier screen that reveals the C-terminal fragment of actin-interacting protein 1 (Aip1) enhances the chemotaxis defect of *Ddnhe1*[−] cells but has no effect in wild-type Ax2 cells. However, expression of full-length Aip1 mostly suppresses chemotaxis defects of *Ddnhe1*[−] cells and restores F-actin assembly. Aip1 functions to promote cofilin-dependent actin remodeling, and we found that although full-length Aip1 binds cofilin and F-actin, the C-terminal fragment binds cofilin but not F-actin. Because pH-dependent cofilin activity is attenuated in mammalian cells lacking H⁺ efflux by NHE1, our current data suggest that full-length Aip1 facilitates F-actin assembly when cofilin activity is limited. We predict the C-terminus of Aip1 enhances defective chemotaxis of *Ddnhe1*[−] cells by sequestering the limited amount of active cofilin without promoting F-actin assembly. Our findings indicate a cooperative role of Aip1 and cofilin in pH-dependent cell migration, and they suggest defective chemotaxis in *Ddnhe1*[−] cells is determined primarily by loss of cofilin-dependent actin dynamics.

INTRODUCTION

In migrating cells, a network of a filamentous actin (F-actin) at the cell front drives membrane protrusion at the leading edge. The dynamic assembly of this actin network in response to migratory cues is tightly regulated by two key molecules; the Arp2/3 complex and cofilin (Pollard and Borisy, 2003). The Arp2/3 complex builds cross-linked actin arrays by nucleating new filaments from the sides of pre-existing filaments (Goley and Welch, 2006). In mammalian cells, cofilin plays an essential role in F-actin assembly by severing filaments to generate new free barbed (plus) ends for nucleation by the Arp2/3 complex (Mouneimne *et al.*, 2006). Cofilin activity is stimulated by dephosphorylation, dissociation from phosphoinositides in the plasma membrane, and an increase in intracellular pH (van Rheenen *et al.*, 2007; Frantz *et al.*, 2008; Bernstein and Bamberg, 2010). In addition to these direct regulatory mechanisms, cofilin-dependent remodeling of F-actin is enhanced by its interaction with actin-interacting protein 1 (Aip1) (Ono, 2003; Briehar *et*

al., 2006). However, how Aip1 functions with regulated cofilin activity in cells is unclear.

Aip1 regulates F-actin dynamics only in the presence of cofilin (Okada *et al.*, 1999; Mohri *et al.*, 2004, 2006; Ono *et al.*, 2004). A member of the WD repeat family with two β -propeller domains, Aip1 forms a tertiary structure with cofilin and F-actin (Voegtli *et al.*, 2003) and lowers the critical concentration of cofilin necessary for F-actin remodeling (Briehar *et al.*, 2006). Aip1 also is reported to cap barbed ends of cofilin-severed filaments (Okada *et al.*, 2002; Balcer *et al.*, 2003), although this function is controversial (Clark *et al.*, 2006; Okada *et al.*, 2006). Loss-of-function studies suggest that Aip1 is required for correct remodeling of F-actin. In *Saccharomyces cerevisiae*, an Aip1-null mutant is a synthetic lethal with mutant cofilin alleles and has thickened actin cables caused by cofilin mislocalization (Rodal *et al.*, 1999). Depletion of UNC-78, an Aip1 homologue in *Caenorhabditis elegans*, results in disorganization of F-actin in the body wall muscle (Ono, 2001). Knockdown of Aip1 in mammalian cells leads to aberrant cytokinesis and inhibits directed migration (Li *et al.*, 2007; Kato *et al.*, 2008). Also in the motile amoebae *Dictyostelium discoideum*, Aip1-null cells have impaired actin-dependent processes, including phagocytosis, cytokinesis, and motility (Konzok *et al.*, 1999).

A rapid assembly of F-actin at the leading edge is necessary for chemotaxis of amoeboid cells such as leukocytes and *Dictyostelium* (Parent, 2004). *Dictyostelium* is an important model for studying chemotactic migration because the mechanics and regulation of F-actin dynamics are similar to those in migrating mammalian cells (Sasaki and Firtel, 2006). In response to the chemoattractant cAMP, *Dictyostelium* cells adopt a polarized, elongated morphology with an F-actin network enriched at the leading edge. Evidence in *Dictyostelium* (Van Duijn and Inouye, 1991; Patel and Barber, 2005)

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Address correspondence to: Diane L. Barber (diane.barber@ucsf.edu).

Abbreviations used: Aip1, actin-interacting protein 1; Dd, *Dictyostelium discoideum*; NHE1, Na⁺-H⁺ exchanger isoform 1.

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and mammalian (Denker and Barber, 2002; Paradiso *et al.*, 2004; Stock and Schwab, 2006) cells indicates that an increase in intracellular pH (pH_i) is necessary for directed migration and for de novo assembly of F-actin at the cell front. *Dictyostelium* cells null for a Na⁺-H⁺ exchanger (*Ddnhe1*[−]) that regulates dynamic changes in pH_i lack efficient chemotaxis and have decreased abundance of F-actin in response to cAMP.

To further understand how *DdNHE1* regulates chemotaxis we used an overexpression library to screen for modifiers of the *Ddnhe1*[−] phenotype. One clone that enhanced *Ddnhe1*[−] chemotaxis and lacked de novo F-actin assembly in response to cAMP contained a C-terminal fragment of *DdAip1*. However, expression of full-length wild-type but not inactive *DdAip1* in *Ddnhe1*[−] cells suppressed the defective chemotaxis phenotype and restored F-actin abundance. Because cofilin-dependent F-actin remodeling in migrating mammalian cells requires increased NHE1 activity and pH_i (Frantz *et al.*, 2008), our findings suggest a cooperative role of Aip1 and cofilin in pH-dependent cell migration.

MATERIALS AND METHODS

Strain, Cell Culture, and Development

Wild-type Ax2 and *Ddnhe1*[−] cells (Patel and Barber, 2005) were cultured in axenic HL5 medium supplemented with 100 U/ml penicillin and 100 µg/ml streptomycin. Plasmid constructs were introduced into Ax2 and *Ddnhe1*[−] cells by electroporation (Knecht and Pang, 1995), and cells expressing *DdAip1* constructs were selected in HL5 medium containing 10 µg/ml G418. Cells expressing Lifeact-monomeric red fluorescent protein (mRFP) were selected with 20 µg/ml hygromycin. For submerged development, exponentially growing cells were washed twice with PB buffer (20 mM K₂HPO₄/KH₂PO₄, pH 6.8) and were allowed to develop at a density of 1 × 10⁶ cells/cm² in PB buffer. Time-lapse images were acquired at 15-min intervals for 20 h by using an Axiovert S-100 microscope (Carl Zeiss, Jena, Germany).

Library Screening

An overexpression cDNA library (provided by Douglas Robinson, Johns Hopkins University, Baltimore, MD) described previously (Robinson and Spudich, 2000) was amplified and electroporated into *Ddnhe1*[−] cells. pREP, a helper plasmid containing an open reading frame (ORF) was cotransformed because these cells have no ORF required for the replication of Ddp2-based pLD1A15SN used for library construction (Robinson and Spudich, 2000). Transformed cells were selected by resistance to 10 µg/ml G418 for 3–4 d, plated onto PB buffer agar plates with a suspension of heat-killed bacteria, and plaque size was scored. Plasmids were recovered from cells using the DNA Mini-prep kit (QIAGEN, Valencia, CA), and cDNA in the plasmids was amplified by polymerase chain reaction (PCR) by using primers LD15M, 5'-AAAAGTCGACCCACGCGTCC-3' and LD13, 5'-CGCGTTTATTATTAGCGGCGGCC-3' and then subcloned into pCR2.1-TOPO vector (Invitrogen, Carlsbad, CA). Purification from *Escherichia coli* DH5α and DNA sequencing revealed a cDNA fragment of *DdAip1* in one clone. The developmental morphology of selected clones was observed under submerged conditions and expression of developmental markers at 0 and 6 h after development was determined by reverse transcriptase-PCR using primers for cAR1 (forward, 5'-GTAATTACATATGGTAG-3' and reverse, 5'-GAAATTAATGGTAAAC-3') and for Ga2 (forward, 5'-GTGTCAAACAG-GCAATG-3' and reverse, 5'-GTCATAACACGAGTATG-3'). *TalA* expression, an internal control, was determined using primers GGATCCATGGATTA-CAAAAAAACATAGACC (forward) and GACGTCGTATGGGTAAAT-GATGTCATACC (reverse).

Plasmid Construction

The plasmid containing full-length of *DdAip1* tagged with green fluorescent protein (GFP) (*DdAip1*-FL; a generous gift from Annette Muller-Taubenberg, Ludwig Maximilians University, Munich, Germany) was used for expression in *Dictyostelium* and as a template for PCR to generate mutant *DdAip1*. The expression vector for *DdAip1*-Δ382 was constructed by PCR amplification using primers AIP1-S, 5'-GGATCCATGGATGATAGTGTTA-3' and Aip1-R2, 5'-GGATCCTTAATTTGATACATACCA-3' and ligation into a BamHI site of pTX-FLAG vector. To express an F-actin reporter in cells, the Lifeact-mRFP sequence (Riedl *et al.*, 2008) was amplified by PCR using primers Lifeact-F, 5'-AAAAGATCTAAAAATGGGTGTGCGAGATTTG-3' and Lifeact-R2, 5'-TTTCTCGAGTTAAGCGCCTGTGCTATG-3' and cloned into BglIII and XhoI sites of EXP5(+) vector containing a hygromycin-resistant cassette. For cloning of rat Aip1, cDNA was synthesized using total RNA from MTLn3 rat adenocarcinoma cells and was used as a template for PCR. Aip1-FL

was amplified by PCR using primers RnAip1-F, 5'-GGATCCATGCCGTAC-GAGATCAA-3' and RnAip1-R, 5'-CTCGAGTTTCAGTAGGTFATTGT-3' and Aip1-Δ382 was amplified using primers RnAip1-TF, ATGGACGACACAGT-GCGGTATACT-3' and RnAip1-R. For production of recombinant proteins, PCR products were cloned into BamHI and XhoI sites of pGEX6P-2 vector (GE healthcare, Little Chalfont, Buckinghamshire, United Kingdom). *DdAip1* with alanine substitutions of E125, E167, F181, and F193 (*DdAip1*-4X) was generated using a QuikChange site-directed mutagenesis kit (Stratagene, La Jolla, CA) using these forward primers (Daip1E125A-S, 5'-GTTGGTGAT-GGTAAAGCTAGATTGGTGCAGCC-3'; Daip1E167A-S, 5'-GCCGCCACT-GGTAGTGCTGATTTGTCAGTC-3'; Daip1F181A-S, 5'-GGTCCACCATTCAAAGCTCAAAAGAATATTGCC-3'; and Daip1F193A-S, 5'-GGTGATT-TCACTCGTGCTGTAATTTGTGTTAG-3').

Chemotaxis Assay

To induce chemotaxis competence, cells were suspended at a density of 1 × 10⁷ cells/ml in PB buffer and pulsed with 30 nM cAMP every 6 min for 5 h (Mendoza and Firtel, 2006). For chemotaxis assays, aliquots of 5-h developed cells were harvested and transferred to 30-mm dishes. cAMP (100 µM) was delivered by a micropipette, and cell movement was recorded at 6-s intervals for 30 min by using an Axiovert S-100 microscope (Carl Zeiss). Cell movement was tracked using ImageJ software (National Institutes of Health, Bethesda, MD), with a manual tracking plug-in. Chemotaxis parameters were calculated by Prism software (GraphPad Software, San Diego, CA), and statistical significance was determined by unpaired two-tailed *t* test.

Actin Polymerization Assay

Abundance of F-actin was determined using two methods. To test clones from the library screen we used Coomassie staining of Triton X-100-insoluble fractions as described previously (Patel and Barber, 2005). To test cells expressing recombinant Aip1 we measured fluorescence of lysates prepared from cells labeled with rhodamine phalloidin as described previously (Chen *et al.*, 2003). Cells pulsed with cAMP for 5 h were resuspended at a density of 3 × 10⁷ cells/ml in PB buffer and incubated with 5 mM caffeine for 20 min to inhibit endogenous cAMP production. After being exposed to a uniform concentration of 2 µM exogenous cAMP, 3 × 10⁶ cells were removed at the indicated times, fixed with 3.7% formaldehyde, and incubated in buffer containing 10 mM PIPES, 0.2% Triton X-100, 20 mM K₂HPO₄/KH₂PO₄, 5 mM EGTA, 2 mM MgCl₂, and 0.4 U of rhodamine-phalloidin (Invitrogen) for 1 h. A Triton X-100-insoluble fraction was obtained by centrifugation and rhodamine-phalloidin bound to F-actin was eluted in methanol overnight. The fluorescence intensity of rhodamine-phalloidin was measured using a SpectraMax M5 plate reader (GE Healthcare).

F-actin Localization

F-actin localization was determined using two approaches. For live cell imaging, cells expressing Lifeact-mRFP were developed by pulsing with cAMP for 5 h and then loaded onto Dunn chemotaxis chambers (Hawksley Technology, Lancing, United Kingdom) in the presence of a cAMP gradient (0–5 µM). Fluorescence images were acquired every 5 s for 10 min with a 60× numerical aperture 1.20 objective lens (Plan Apo; Nikon, Tokyo, Japan) on a TE2000 inverted microscope (Nikon) equipped with a spinning-disk unit (CSU10; Yokogawa Electronics, Tokyo, Japan), 488-nm and 561-nm lasers, and a CoolSNAP HQ2 camera (Photometrics, Tucson, AZ). We also used phalloidin labeling of fixed cells chemotaxing toward cAMP released from a pipette, as described for chemotaxis assays. For analysis of F-actin localization, fluorescence intensity of rhodamine-phalloidin was measured along the cell perimeter using ImageJ. Fluorescence intensity in arbitrary units in 20 sectors around the cell starting from the cell rear was determined.

In Vitro Aip1 and Cofilin Binding

Recombinant wild-type and mutant rat Aip1 expressed as a glutathione-S-transferase-tagged fusion in *E. coli* BL21 (DE3) was purified using glutathione-Sepharose 4B (GE Healthcare) according to manufacturer's instructions. Recombinant wild-type cofilin was expressed and purified as described previously (Frantz *et al.*, 2008). For binding assays, 20 µM cofilin was incubated for 1 h with 0.2 µM GST or GST-Aip1 bound to glutathione-Sepharose beads in incubation buffer (20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl₂, and 0.1% NP40). After washing with incubation buffer, unbound cofilin was removed and beads-bound cofilin was boiled in the SDS-sample buffer and resolved by SDS-polyacrylamide gel electrophoresis (PAGE).

Actin Cosedimentation Assay

Rabbit muscle actin (Cytoskeleton, Denver, CO) was polymerized in 50 mM KCl, 2 mM MgCl₂, and 1 mM ATP, and 10 µg was incubated with 0.5 µM GST and GST fusion proteins for 30 min at 24°C. F-actin was pelleted by centrifugation at 100,000 × *g* for 20 min, and proteins in supernatant and pellet fractions were resolved by 12% SDS-PAGE.

Immunoblotting

Exponentially growing cells were lysed by sonicating in lysis buffer (1× phosphate-buffered saline, protease inhibitors) and proteins in total cell lysates were separated by SDS-PAGE and transferred to polyvinylidene difluoride membranes. Membranes were probed with antibodies for GFP (JL-8, 1:1000; Clontech, Mountain View, CA) and β -actin (C4, 1:5000; Millipore Bioscience Research Reagents, Temecula, CA), and bound antibody was detected by enhanced chemiluminescence (Pierce Chemical, Rockford, IL).

RESULTS

A C-Terminal Fragment of *DdAip1* Enhances Impaired Chemotaxis of *Ddnhe1*[−] Cells

To determine how *DdNHE1* regulates chemotaxis of *Dictyostelium* cells, we developed an easily scored assay to test for phenotypic changes. Although expression of *DdNHE1* is very low in vegetative cells, we found that *Ddnhe1*[−] cells produced substantially smaller plaques than wild-type Ax2 cells on a heat-killed bacterial lawn. We screened a cDNA overexpression library previously designed to identify suppressors of cytokinesis-defective mutants (Robinson and Spudich, 2000) and scored for plaque formation. Although no clones were identified with a suppressed phenotype, one clone (clone 15) had an enhanced phenotype in which the plaques were smaller than those produced by *DdNhe1*[−] cells. We analyzed the development of clone 15 by determining stream formation, which is an index of efficient chemotaxis. When starved under nonnutrient phosphate buffer, Ax2 cells formed streams at 8 h and aggregates at 24 h (Figure 1A). *Ddnhe1*[−] cells had delayed stream formation.

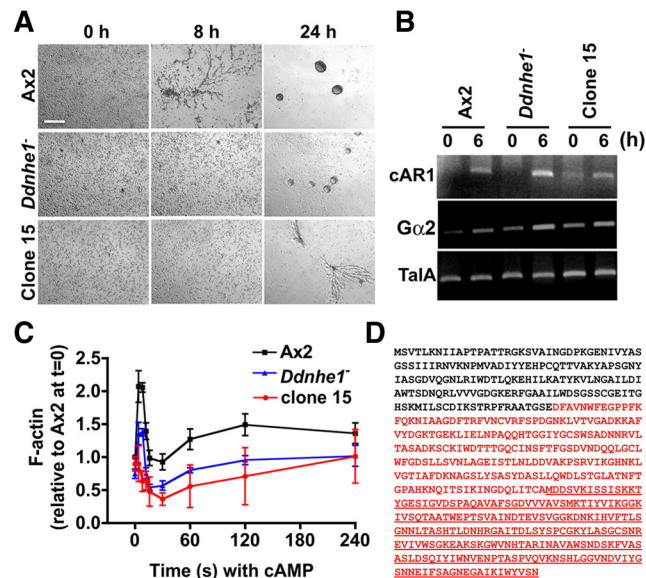


Figure 1. Modifier screen reveals the C-terminal fragment of Aip1 enhances defects in development and F-actin assembly in *Ddnhe1*[−] cells. (A) Developmental morphologies under nonnutrient buffer at the indicated times. Clone 15 cells formed aggregates more slowly than parental *Ddnhe1*[−] cells. Bar, 100 μ m. (B) Expression of early developmental genes *cAR1* and *Gα2* determined by RT-PCR was similar in clone 15 and Ax2 cells. *TalA* expression was used as a loading control. (C) Time-dependent increase in total F-actin in response to cAMP was abolished in clone 15 cells. Data are expressed as the means \pm SEM of three independent experiments. (D) Amino acid sequence of *DdAip1*. The sequence in red denotes a truncated *DdAip1* sequence recovered from clone 15. The underlined sequence denotes *DdAip1*- Δ 382 predicted from a methionine start site.

Although aggregates were seen at 24 h, they were smaller than those with Ax2 cells. Development of clone 15 was more delayed than *Ddnhe1*[−] cells (Figure 1A), with stream formation but no aggregates at 24 h. Development and stream formation require induced expression of several aggregation genes, including the G protein-coupled cAMP receptor *cAR1* and the G protein subunit *Gα2*. Expression of these genes at 6 h of development was similar in the three strains, indicating that starvation-induced gene expression was unaffected (Figure 1B).

In response to chemotactic cues, *Dictyostelium* cells have a well-characterized biphasic increase in F-actin, including a rapid and transient first peak at 4–8 s and a slower and more prolonged second peak at 30–180 s (Condeelis *et al.*, 1988; Chen *et al.*, 2003). We showed previously that in *Ddnhe1*[−] cells the first peak is reduced by 50% and the second peak is largely absent (Patel and Barber, 2005). Clone 15 had markedly less de novo increases in F-actin. Most notably, there was no first peak, which further indicated the *Ddnhe1*[−] phenotype was enhanced (Figure 1C).

Sequencing of the recovered plasmid from clone 15 revealed that it contained a C-terminal fragment of a gene encoding *DdAip1* (Figure 1D). Because the recovered cDNA did not include the full-length *DdAip1* sequence but contained a methionine residue at position 382, we predicted it encoded a *DdAip1*- Δ 382 fragment (Figure 1D). This finding was revealing with regard to the role of Aip1 in promoting cofilin-dependent actin dynamics (Ono, 2003), and our previous data showing that NHE1 activity and increased pH_i are necessary for cofilin-mediated F-actin remodeling in migrating mammalian fibroblasts (Frantz *et al.*, 2008).

Chemotaxis Defect of *Ddnhe1*[−] Cells Is Enhanced by *DdAip1*- Δ 382 but Suppressed by Full-Length *DdAip1*

To confirm that *DdAip1*- Δ 382 enhances the phenotype of *Ddnhe1*[−] cells, *DdAip1*- Δ 382 tagged with a FLAG epitope was transformed into Ax2 (Ax2/*DdAip1*- Δ 382) and *Ddnhe1*[−] (*Ddnhe1*[−]/*DdAip1*- Δ 382) cells. We showed previously that the expression of *Gα2* and *cAR1* in *Ddnhe1*[−] cells is similar between 3 and 6 h (Patel and Barber, 2005), and hence determined chemotaxis toward a point source of cAMP in cells starved for 5 h. Ax2 cells rapidly adopted a polarized morphology with leading edge membrane protrusions oriented toward cAMP (Figure 2, A and C, and Supplemental Video 1). They migrated efficiently with a speed of 13.09 ± 0.23 μ m/min (Figure 2D) and a chemotactic index of 0.95 ± 0.010 (Figure 2E). *Ddnhe1*[−] cells failed to adopt a polarized morphology, and although they did extend membrane protrusions, these were not directed toward the micropipette (Figure 2, A and C, and Supplemental Video 2). In addition, *Ddnhe1*[−] cells moved randomly and more slowly with a speed of 4.54 ± 0.36 μ m/min (Figure 2D) and a chemotactic index of 0.63 ± 0.045 (Figure 2E). *Ddnhe1*[−]/*DdAip1*- Δ 382 cells remained rounded without visible membrane protrusions and showed limited movement (Figure 2, A and C, and Supplemental Video 4). Their speed of 2.67 ± 0.24 μ m/min (Figure 2D) and chemotactic index of 0.39 ± 0.045 (Figure 2E) were significantly less than those of *Ddnhe1*[−] cells ($p = 0.0001$, $n = 20$), suggesting a more defective chemotaxis. In contrast, expression of *DdAip1*- Δ 382 in Ax2 cells (Ax2/*DdAip1*- Δ 382) had no effect on chemotaxis, including polarity (Figure 2, A and C, and Supplemental Video 3), speed (Figure 2D), and chemotactic index (Figure 2E). Immunoblotting confirmed comparable expression of *DdAip1*- Δ 382 in Ax2 and *Ddnhe1*[−] cells (Supplemental Figure 2A). These findings and our screening data suggest that *DdAip1*- Δ 382

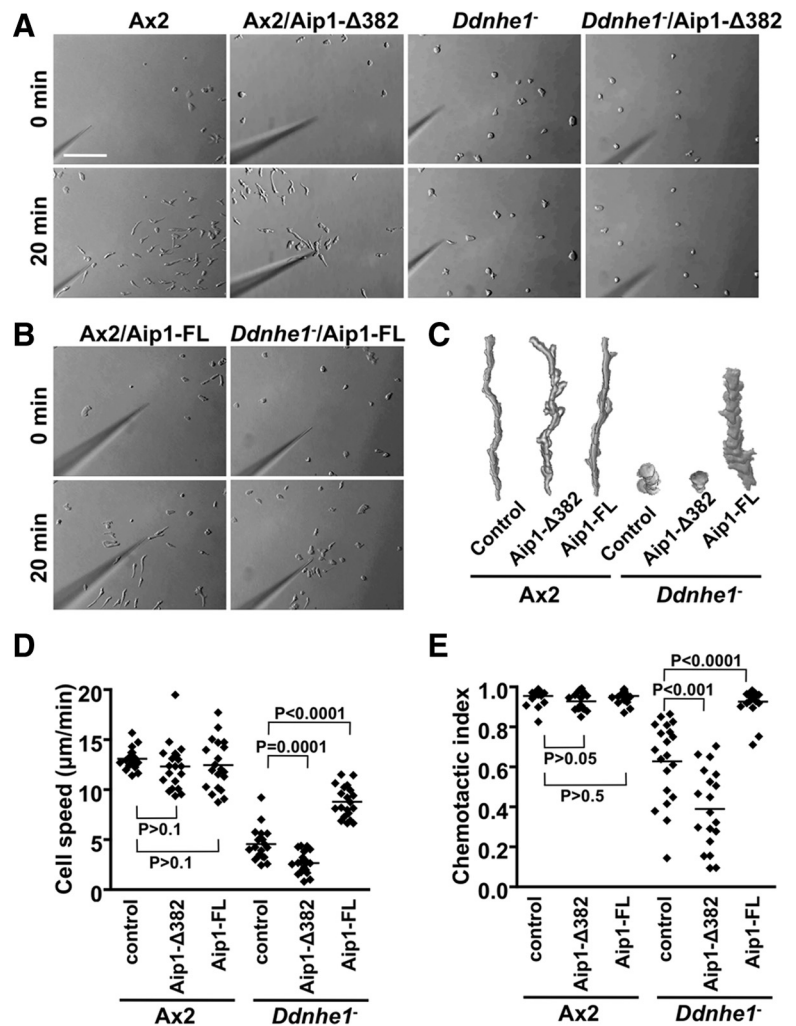


Figure 2. Impaired chemotaxis of *Ddnhe1*⁻ cells is enhanced by *DdAip1*-Δ382 but suppressed by *DdAip1*-FL. Chemotactic movement toward a micropipette tip filled with cAMP was recorded for 30 min. (A) Images of Ax2 and *Ddnhe1*⁻ cells without and with heterologous expression of *DdAip1*-Δ382. (B) Images of Ax2 and *Ddnhe1*⁻ cells expressing *DdAip1*-FL. (C) Morphology and tracking of the indicated cell types determined by drawing and overlapping images of a single cell from frames taken at 1-min intervals. (D) Speed of the indicated cell types, expressed as total distance moved divided by total moving time. The speed of *Ddnhe1*⁻ cells is significantly decreased by *DdAip1*-Δ382 but increased by *DdAip1*-FL. (E) chemotactic index of the indicated cell types, determined as net distance moved divided by total moving distance during the time period. The chemotactic index of *Ddnhe1*⁻ cells is significantly decreased by *DdAip1*-Δ382 but increased by *DdAip1*-FL. Each dot in the scatter plots represents an individual cell in time-lapse images, and data are representative of three independent experiments.

enhances the impaired chemotaxis phenotype of *Ddnhe1*⁻ cells.

Having confirmed that *DdAip1*-Δ382 enhanced *Ddnhe1*⁻ chemotaxis, we next determined the effect of expressing full-length *DdAip1* (*DdAip1*-FL) in Ax2 and *Ddnhe1*⁻ cells (Ax2/*DdAip1*-FL and *Ddnhe1*⁻/*DdAip1*-FL, respectively). In contrast to *DdAip1*-Δ382, impaired chemotaxis of *Ddnhe1*⁻ cells was mostly restored by expression of *DdAip1*-FL. *Ddnhe1*⁻/*DdAip1*-FL cells adopted a polarized morphology, although they were less elongated than Ax2 cells (Figure 2, B and C, and Supplemental Video 6). Their speed ($8.79 \pm 0.35 \mu\text{m}/\text{min}$) and chemotactic index (0.93 ± 0.016) were 67 and 98%, respectively, of Ax2 cells. (Figure 2, D and E). For Ax2/*DdAip1*-FL cells, migration speed (Figure 2D; $p > 0.5$, $n = 20$; Supplemental Video 5) and chemotactic index (Figure 2E; $p > 0.05$, $n = 15$) were not significantly different from Ax2 cells. Comparable expression of *DdAip1*-FL in Ax2 and *Ddnhe1*⁻ cells was confirmed by immunoblotting (Supplemental Figure 2B). These data indicate that in contrast to *DdAip1*-Δ382, full length of *DdAip1* suppresses impaired chemotaxis of *Ddnhe1*⁻ cells.

Because our library screen did not identify *DdAip1*-FL as a modifier of the *Ddnhe1*⁻ phenotype, we tested plaque formation on heat-killed bacteria by cells transformed with *DdAip1*-FL and *DdAip1*-Δ382. After 2 d, Ax2 cells formed large plaques with fruiting bodies, whereas *Ddnhe1*⁻ cells

formed smaller plaques and did not aggregate until 2 d (Supplemental Figure 1). As expected, *Ddnhe1*⁻/*DdAip1*-Δ382 cells formed smaller plaques than *Ddnhe1*⁻ cells, whereas *Ddnhe1*⁻/*DdAip1*-FL cells formed larger plaques with some aggregates compared with *Ddnhe1*⁻ cells (Supplemental Figure 1). These data indicate that expression of *DdAip1*-FL partially rescues defective plaque formation by *Ddnhe1*⁻ cells. Because the library we screened contains inserts with an average size of 1.1–1.3 kb (Robinson and Spudich, 2000), we suspect that the 1.8-kb *DdAip1*-FL might not be expressed or is expressed in low abundance.

DdAip1-FL Restores F-actin Assembly but Not Polarity of *Ddnhe1*⁻ Cells

Because clone 15 had markedly attenuated F-actin with cAMP, we measured the kinetics of F-actin in response to uniform cAMP. Ax2 and Ax2/*DdAip1*-Δ382 cells had a similar amount of F-actin in the first and second phases (Figure 3A). However, as with clone 15, *Ddnhe1*⁻/*DdAip1*-Δ382 cells showed a markedly attenuated peak of F-actin in both first and second phases (Figures 1B and 3A). In contrast to *Ddnhe1*⁻/*DdAip1*-Δ382 cells, *Ddnhe1*⁻/*DdAip1*-FL cells showed restored F-actin abundance in both first and second phases to levels seen with Ax2 cells (Figure 3B). The kinetics and abundance of F-actin in Ax2/*DdAip1*-FL cells were similar to Ax2 cells (Figure 3B). Of importance, all strains had a

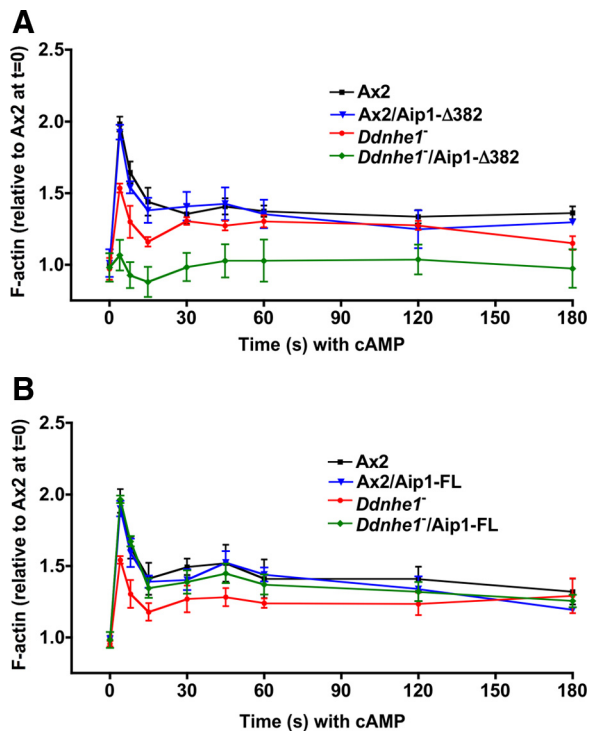


Figure 3. Stimulated F-actin assembly in *Ddnhe1*[−] cells is attenuated by *DdAip1*-Δ382 but rescued by *DdAip1*-FL. Total F-actin was determined by fluorescence of rhodamine-phalloidin in Triton X-100-insoluble fractions of cells at the indicated times (seconds) after addition of cAMP and expressed relative to F-actin of Ax2 at time 0 in the absence of cAMP. (A) Expression of *DdAip1*-Δ382 markedly attenuated the first and second peaks of F-actin in *Ddnhe1*[−] cells but had no effect in Ax2 cells. (B) Expression of *DdAip1*-FL in *Ddnhe1*[−] cells restores abundance of F-actin in the first peak to that in Ax2 cells.

similar abundance of F-actin in the absence of cAMP (0 time), which confirms our previous findings in *Dictyostelium* (Patel and Barber, 2005) and mammalian (Frantz *et al.*, 2008) cells that NHE1 activity is necessary for de novo assembly of F-actin in response to chemoattractants but not for steady-state amounts of F-actin.

Although *DdAip1*-FL restored chemotaxis and F-actin assembly in *Ddnhe1*[−] cells, it only partially rescued polarity. Morphological polarity was scored by roundness, which is an index for lack of elongated shape and front-back asymmetry of *Dictyostelium* cells (van Es *et al.*, 2001). A higher percentage of roundness indicates less polarized cells. Starved Ax2 cells were mostly elongated, had a roundness index of $47 \pm 2.3\%$ and extended pseudopods at the front toward the point source of cAMP (Figure 4, A and B). In contrast, *Ddnhe1*[−] cells had a higher roundness index of $71 \pm 1.4\%$, and they extended smaller pseudopods that were not restricted to the cell front. The morphology of Ax2/*DdAip1*-Δ382 and Ax2/*DdAip1*-FL was similar to that of Ax2 cells. *Ddnhe1*[−]/*DdAip1*-Δ382 cells showed increased roundness of $78 \pm 2.1\%$ compared with *Ddnhe1*[−] cells (Figure 4B). *Ddnhe1*[−]/*DdAip1*-FL cells were more polarized than *Ddnhe1*[−] cells with a significantly decreased roundness index of $59 \pm 3.3\%$ compared with *Ddnhe1*[−] cells. However, *Ddnhe1*[−]/*DdAip1*-FL cells were still less polarized than Ax2 ($47 \pm 2.3\%$) or Ax2/*DdAip1*-FL cells ($45 \pm 2.8\%$) (Figure 4, A and B). We also determined F-actin localization in migrating cells as another index of cell polarity. Cells transformed

with the F-actin reporter Lifeact (Riedl *et al.*, 2008) were starved for 5 h, transferred to Dunn chemotaxis chambers, and Lifeact-mRFP fluorescence was imaged in cells migrating across a cAMP gradient. Fluorescence, an index of F-actin, was restricted to pseudopods at the leading edge of migrating Ax2 cells but was localized in multiple protrusions around the cortex of *Ddnhe1*[−] cells (Figure 4C). In *Ddnhe1*[−]/*DdAip1*-FL cells, fluorescence was seen at the leading edge, but also at lateral edges (Figure 4C). Similar findings were obtained using phalloidin-rhodamine labeling of F-actin (Supplemental Figure 3). Together, these data suggest that although expression of *DdAip1*-FL mostly restores the chemotaxis index of *Ddnhe1*[−] cells, it only partially restores their leading-edge localization of F-actin and their polarity.

Aip1-FL but Not *Aip1*-Δ382 Binds F-actin

The established function of Aip1 is to enhance cofilin severing or disassembly of actin filaments (Ono, 2003). We showed previously that F-actin severing and the generation of new free barbed ends by cofilin are attenuated in migrating fibroblasts lacking NHE1 activity and having low pH_i (Frantz *et al.*, 2008). Hence, our current data suggest that expression of *DdAip1*-FL may restore F-actin assembly in *Ddnhe1*[−] cells by enhancing reduced cofilin activity at low pH_i. However, to understand why the *Ddnhe1*[−] chemotaxis phenotype is suppressed by *DdAip1*-FL but enhanced by *DdAip1*-Δ382 we asked whether these two proteins differed in their ability to bind cofilin and F-actin. Mutagenesis studies indicate that Aip1 residues important for binding to cofilin and F-actin are located in both of N- and C-terminal propellers (Clark *et al.*, 2006; Okada *et al.*, 2006; Clark and Amberg, 2007). Because *Dictyostelium* has six cofilin isoforms and it is unknown which isoforms binds *DdAip1*, we used recombinant rat cofilin and Aip1 to test binding. We found that cofilin bound to GST-tagged Aip1-FL and to Aip1-Δ383 but not to GST alone (Figure 5A). These data are significant because they show that the C terminus of Aip1 is sufficient to bind cofilin in the absence of the N terminus. We tested whether binding is regulated by changes in pH but found that binding was pH-independent between pH 6.8–7.8 (data not shown). Using a sedimentation assay, we found that Aip1-FL but not Aip1-Δ383 copelleted with F-actin (Figure 5B), which indicates that the N-terminal propeller is required for binding to F-actin. These data suggest that Aip1 suppression of the *Ddnhe1*[−] chemotaxis phenotype probably requires its binding to F-actin.

Inactive *DdAip1* Does Not Restore Impaired Chemotaxis or F-actin Assembly of *Ddnhe1*[−] Cells

A mutant “inactive” *C. elegans* Aip1 (UNC-78) containing alanine substitution of four residues (E126, D168, F182, and F192) in the N-terminal β-propeller is unable to enhance cofilin-dependent depolymerization and to suppress an *unc-78*-null phenotype (Mohri *et al.*, 2006). Because these four residues are conserved in *DdAip1* (E125, E167, F181, and F193), we determined effects of a similar mutant (*DdAip1*-4X) tagged at the N terminus with GFP to further understand the difference between *DdAip1*-FL and *DdAip1*-Δ382 on chemotaxis of *Ddnhe1*[−] cells. Immunoblotting for GFP indicated similar expression of *DdAip1*-4X in Ax2 and *Ddnhe1*[−] cells (Supplemental Figure 2B). Like the homologous *C. elegans* mutant, *DdAip1*-4X bound to cofilin and F-actin (Supplemental Figure 4), suggesting that protein structure is retained. Under submerged condition, starved Ax2 cells streamed and formed tight aggregates at 20 h (Figure 6A and Supplemental Video 7). Expression of

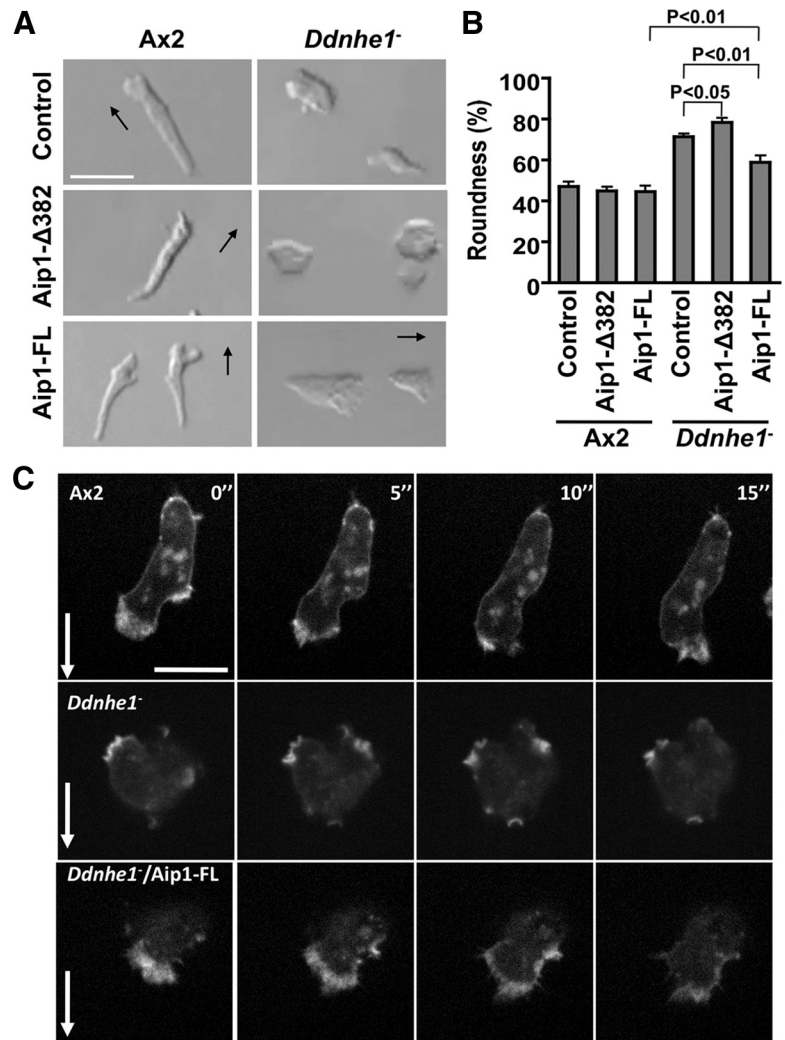


Figure 4. *DdAip1*-FL does not restore impaired polarity of *Ddnhe1*⁻ cells. (A) Morphology of cells migrating toward cAMP. Arrows indicate the direction of cell migration. Bar, 10 μ m. (B) Cell roundness, calculated as $100 \times 4\pi \times \text{cell area}/(\text{cell perimeter})^2$ indicated that *Ddnhe1*⁻ cells were more elongated with expression of *DdAip1*-FL. Data are expressed as the means \pm SEM from 10 cells in three independent time-lapse sequences. (C) Dynamics of F-actin in chemotaxing cells expressing Lifeact-mRFP was determined by live cell fluorescence imaging. Arrows indicate direction of increasing cAMP gradient in the Dunn chemotaxis chamber. Lifeact-mRFP fluorescence is restricted to leading edge pseudopods of Ax2 cells but is seen in numerous protrusions not localized to the direction of the cAMP gradient in *Ddnhe1*⁻ cells. In *Ddnhe1*⁻/*DdAip1*-FL cells, fluorescence is localized at the front and at lateral edges of migrating cells. Bar, 10 μ m.

DdAip1-FL or 4X did not change aggregation timing of Ax2 cells (data not shown). Streaming and aggregation were delayed in *Ddnhe1*⁻ cells but restored with expression of *DdAip1*-FL (Figure 6A and Supplemental Videos 8 and 9). However, expression of *DdAip1*-4X had no effect on streaming and aggregation of *Ddnhe1*⁻ cells (Figure 6A and Supplemental Video 10), indicating the phenotype was not suppressed or enhanced. In addition, chemotaxis (Figure 6B) and F-actin kinetics (Figure 6C) of *Ddnhe1*⁻ cells were similar in the absence and presence of *DdAip1*-4X. These data suggest that the ability of Aip1-FL to restore efficient chemotaxis in *Ddnhe1*⁻ cells requires its activity in enhancing cofilin function.

DISCUSSION

Aip1 is recognized as a cofactor for cofilin and enhances cofilin-dependent F-actin dynamics during endocytosis, cytokinesis, and cell movement. Aip1 facilitates severing and depolymerization of actin filaments only when they are decorated with cofilin (Ono *et al.*, 2004), it lowers the amount of cofilin necessary for F-actin disassembly (Brieher *et al.*, 2006), and it caps free barbed ends of cofilin-severed filaments (Okada *et al.*, 2002; Balcer *et al.*, 2003). Genetic evidence also indicates a functional interaction between Aip1

and cofilin. In yeasts, loss of *aip1* is a synthetic lethal with cofilin mutants and cells have mislocalized cofilin (Rodal *et al.*, 1999). In *C. elegans*, deletion of *unc-78* enhances a motility defect of *unc-60B*, a cofilin homologue, and induces mislocalization of cofilin to actin aggregates (Ono, 2001). In addition, depletion of active cofilin blocks association of Aip1 with actin in *Xenopus* cells (Tsuji *et al.*, 2009), and expression of active cofilin restores cytokinesis and migration that are impaired by knockdown of Aip1 in mammalian cells (Kato *et al.*, 2008). In addition to Aip1, cofilin activity is also enhanced by increased pH (Bernstein and Bamburg, 2004). In motile mammalian fibroblasts, increased severing activity of cofilin for the assembly of new actin filaments requires increased H⁺ efflux by the plasma membrane Na⁺-H⁺ exchanger NHE1 (Frantz *et al.*, 2008). We now show that in *Dictyostelium* cells lacking *Ddnhe1* impaired F-actin assembly and defective chemotaxis are restored by expression of full-length Aip1.

Our findings on a genetic interaction between *DdAip1* and *DdNHE1* suggest that defective chemotaxis in *Ddnhe1*⁻ cells may be determined primarily by loss of cofilin-dependent actin dynamics. A recently recognized function of Aip1 is to lower the critical concentration of active cofilin necessary for F-actin remodeling (Brieher *et al.*, 2006). Hence, we predict that heterologously expressed Aip1-FL increases cofilin ac-

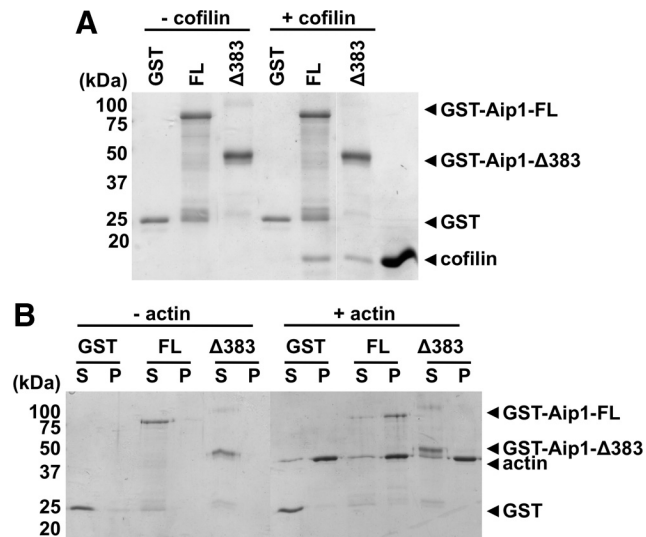


Figure 5. C-Terminal fragment of Aip1 binds cofilin but not F-actin. (A) Binding of GST-Aip1 full length (FL) and C-terminal fragment ($\Delta 383$) and cofilin ($20 \mu\text{M}$) in the absence of actin was determined by GST-pull-down assay. Cofilin binds both Aip1-FL and $\Delta 383$, but not GST alone. (B) Binding of Aip1 to F-actin in the absence of cofilin was determined by using an actin cosedimentation assay. Polymerized actin ($10 \mu\text{M}$) was incubated with $0.5 \mu\text{M}$ Aip1-FL and $\Delta 383$ for 1 h. After centrifugation, the supernatant (S) and the pellet (P) fractions were separated and resolved by SDS-PAGE. Aip1-FL but not $\Delta 383$ binds to actin filaments.

tivity that is likely low in *Ddnhe1*⁻ cells because pH_i is markedly less than in Ax2 cells (Patel and Barber, 2005). In

support of this prediction, expression of Aip1-FL suppresses chemotaxis defects and restores attenuated F-actin assembly of *Ddnhe1*⁻ cells but has no effect in wild-type Ax2 cells. These findings further suggest that the role of *DdAip1* in chemotaxis is a separate but compensatory pathway for cofilin regulation, such as seen with phosphatidylinositol 3-kinase and TORC2 for AKT regulation (Kamimura *et al.*, 2008). Although most studies on Aip1 function describe enhancement of cofilin-mediated F-actin disassembly in vitro, Aip1-FL restores F-actin assembly in *Ddnhe1*⁻ cells, which is consistent with a critical role of cofilin-dependent filament severing in generating new free barbed ends for de novo F-actin assembly (Mouneimne *et al.*, 2006). Okreglak and Drubin (2010) recently showed that the annealing of actin oligomers formed by Aip1 and cofilin is a physiologically relevant pathway for F-actin assembly, suggesting a role of Aip1 and cofilin in actin filament assembly in cells. Biphasic F-actin assembly in response to cAMP is attenuated in *Ddnhe1*⁻ cells (Patel and Barber, 2005), and we found that the first phase is more dramatically rescued by Aip1-FL. Similarly, biphasic formation of free barbed ends in response to growth factors (Mouneimne *et al.*, 2006) is attenuated in mammalian NHE1-deficient cells and only the first phase is restored by expression of a mutant pH-insensitive cofilin (Frantz *et al.*, 2008). Our findings suggest that the first phase of actin assembly in response to cAMP may be cofilin dependent and that Aip1-FL restores attenuated cofilin-dependent actin dynamics in *Ddnhe1*⁻ cells. Because the *Dictyostelium* genome includes six cofilins, it is difficult to know which isoform is specifically enhanced by Aip1 in cells,

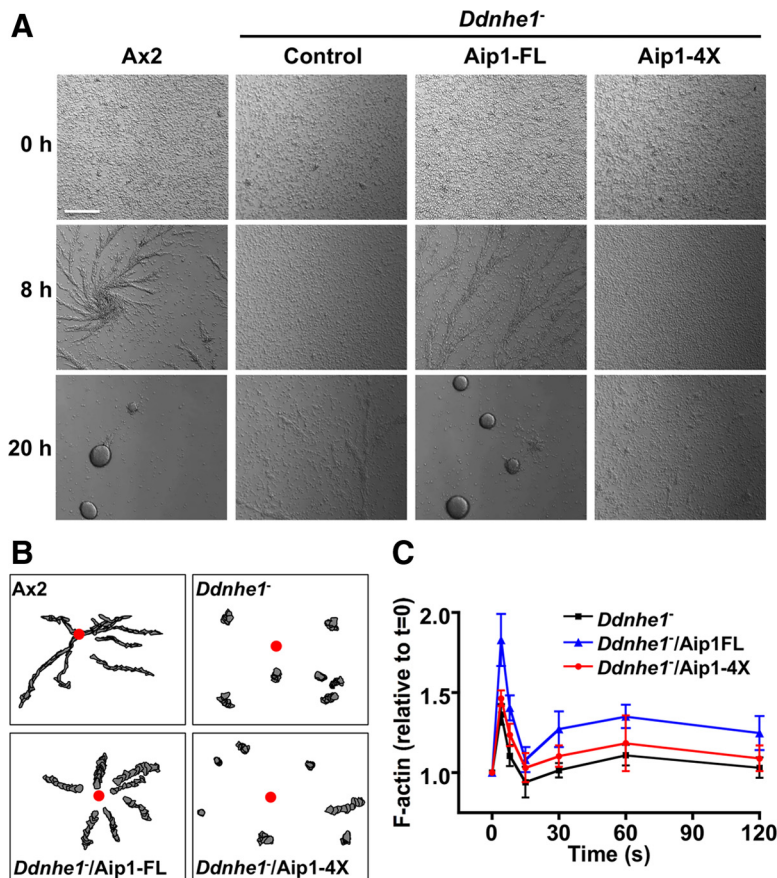


Figure 6. Inactive *DdAip1* does not restore impaired chemotaxis or F-actin assembly of *Ddnhe1*⁻ cells. (A) Analysis of developmental morphology indicated that *Ddnhe1*⁻/*DdAip1*-FL cells form aggregates at 20 h like Ax2 cells, but aggregate formation is not different in *Ddnhe1*⁻/*DdAip1*-4X cells and *Ddnhe1*⁻ cells. (B) *DdAip1*-4X cells did not rescue chemotaxis defect of *Ddnhe1*⁻ cells. The outline of seven cells was drawn and overlapped from the images taken at 1-min intervals. The red dot indicates the position of a micropipette containing cAMP. (C) Time-dependent F-actin amount in response to cAMP showed that expression of *DdAip1*-FL but not *DdAip1*-4X rescued attenuated F-actin assembly of *Ddnhe1*⁻ cells.

although actin disassembly by *Dictyostelium* cofilin 1 is enhanced by Aip1 (Aizawa *et al.*, 1999).

Cofilin dependence for suppression of the *Ddnhe1*[−] phenotype by Aip1-FL is also supported by our findings that Aip1-4X has no effect on impaired chemotaxis and F-actin assembly in *Ddnhe1*[−] cells. In *C. elegans*, a 4X mutant of UNC-78 analogous to Aip1-4X lacks cofilin-dependent actin disassembly and fails to rescue the phenotype of *unc-78*-null mutants (Mohri *et al.*, 2006). Although Aip1 has both severing and capping activity, UNC-78-4X abolishes severing activity but not capping (Mohri *et al.*, 2006). Hence, rescue of the *Ddnhe1*[−] phenotype by Aip1-FL but not Aip1-4X suggests the importance of severing but not capping activity in the absence of *Ddnhe1*. We found that expression of the C-terminal fragment Aip1-Δ382 markedly enhances defects in chemotaxis and F-actin assembly of *Ddnhe1*[−] cells but has no effect on Ax2 cells. Thus, we suspect that Aip1-Δ382 acts as a partial dominant negative, with effects seen when cofilin activity is limited in *Ddnhe1*[−] cells but not in Ax2 cells. We initially predicted this dominant negative effect might reflect increased F-actin capping in cells with limited filament severing activity of cofilin because of decreased p*H*_i. However, although Aip1-Δ382 retains binding to cofilin it is unable to bind F-actin. A more likely interpretation of why Aip1-Δ382 enhances defects in chemotaxis and F-actin assembly of *Ddnhe1*[−] cells is that Aip1-Δ382 may sequester the limited amount of active cofilin in cells with low p*H*_i or prevent free cofilin from binding to F-actin. However, Aip1-4X has little effect in *Ddnhe1*[−] cells, although it has no activity but still binds cofilin. We speculate that in contrast to Aip1-Δ382, the 4X mutant preferentially complexes with F-actin-bound cofilin, consistent with previous findings that the 4X mutation does not disrupt cofilin binding to F-actin but inhibits filament severing by cofilin (Mohri *et al.*, 2006). Because the mechanism whereby Aip1 promotes cofilin activity is still unknown, it is difficult to speculate on the functional significance of these differences between Aip1-Δ382 and Aip1-4X.

Although Aip1-FL restores attenuated F-actin kinetics in *Ddnhe1*[−] cells, it does not completely rescue cell polarity. During chemotaxis, *Ddnhe1*[−] cells expressing Aip1-FL are less elongated than Ax2 cells and actin-rich pseudopods are not restricted to the leading edge, which probably contributes to their slower speed compared with Ax2 cells. In response to chemoattractant, the first phase of actin polymerization in *Dictyostelium* cells is predicted to be important for establishing morphological asymmetry and the second phase drives membrane protrusion (Chen *et al.*, 2003). However, the kinetics and magnitudes of both F-actin phases are similar in Ax2 cells and in *Ddnhe1*[−] cells expressing Aip1-FL, but expression of Aip1-FL does not limit F-actin assembly to the leading edge. Local activation of cofilin at the leading edge of migrating fibroblast cells is necessary to maintain cell polarity during directional migration (Dawe *et al.*, 2003), which may explain why *Ddnhe1*[−] cells are more elongated with Aip1-FL and more rounded with Aip1-Δ382. In motile mammalian fibroblasts, H⁺ efflux by NHE1 is necessary for directional polarity (Denker and Barber, 2002) and forms a bistable positive feedback loop with the low molecular weight GTPase Cdc42 (Frantz *et al.*, 2007). Despite the critical role of Cdc42 in cell polarity of mammalian cells, its *Dictyostelium* orthologue has not been identified and how H⁺ efflux determines polarity in *Dictyostelium* cells is unresolved. Because cofilin localization in protrusions is retained in *Ddaip1*-null cells (Konzok *et al.*, 1999), our data suggest that Aip1 does not function to restrict F-actin assembly at the cell front.

Increased p*H*_i is an evolutionarily conserved signal necessary for directed cell migration. The ability of Aip1-FL to nearly restore impaired chemotaxis in *Ddnhe1*[−] cells highlights the critical importance of pH-dependent cofilin activity that is likely not limited to *Dictyostelium* cells. Cofilin-dependent F-actin remodeling and regulatory pathways promote migration of invasive cancer cells (Wang *et al.*, 2007), and Aip1 also enhances directed migration of mammalian cells (Li *et al.*, 2007; Kato *et al.*, 2008). Moreover, increased p*H*_i is a hallmark of most cancer cells (Cardone *et al.*, 2005; Harguindeguy *et al.*, 2005). Hence, our findings on NHE1 and Aip1 in amoeboid cell migration reveal new insights in understanding the mechanism of cofilin-dependent invasive migration of cancer cells.

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